

Accelerated weathering of cementitious matrix for the development of an accelerated laboratory test of biodeterioration

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Abstract

Cement based materials are porous, may contain organic adjuvants, and thus possesses an important primary bioreceptivity. To preserve constructions from fungal colonization and to act efficiently against fungal biodeterioration, it is necessary to have a better understanding of biodeterioration mechanisms and its effects on materials properties. An accelerated laboratory test which allows us to compare the growth of three fungal strains and the aesthetic biodeterioration of a cementitious matrix was developed. As the surface pH of the fresh cement specimen is too high to allow fungal growth (pH ~12), accelerating weathering of the matrix, consisting of the combination of carbonation and leaching, was performed to reduce the matrix alkalinity. XRD analyses and SEM observations pointed out that the matrix surface is progressively covered by a calcium carbonate layer as the weathering increases. Results point out that the microbial growth occurs on matrix with a surface composition more like a limestone than a cementitious one.

Keywords:

Biodeterioration ; Cement paste ; Carbonation ; Leaching ; Accelerated weathering ; Microbial growth

I. Introduction

Micro-organisms—bacteria, cyanobacteria, fungi, algae, and lichens—are liable to grow on building materials. Biological activity contributes to deterioration of building material, and its interaction with physico-chemical mechanisms is considered central to understanding the long term deterioration [47]. Physical, chemical, and biological agents act in co-association, ranging from synergistic to antagonistic, to deteriorate stone [63].

In Civil Engineering the most widely used material is concrete for different kind of application: bridge construction, sewer pipes, buildings and in some case during restoration of Cultural Heritage Monuments. Cement plays an essential role in concrete works behaviour, because it provides its mechanical resistance [32]. Cement based materials are porous, may contain organic adjuvants, and thus possess an important primary bioreceptivity.

Bioreceptivity, as defined by Guillitte [31], is the totality of materials properties that contribute to the establishment, anchorage and development of fauna and/or flora. Primary bioreceptivity is the initial potential of colonisation [31].

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Fungi are among the most harmful organisms associated to biodeterioration of organic and inorganic materials [58]. Their occurrence on the stones is reported to be combined not only with aesthetic spoiling of the monuments, due to colour changes and black spots, but also there is strong evidence that these organisms can colonize deeper cracks, cause crater shaped lesions, chipping and exfoliation of the rock surface combined with the loss of materials [58, 65].

In general, two main groups of fungi are usually isolated from rock surfaces [8]: (i) one group includes species of the genera of *Hyphomycetes* and *Coelomycetes* among which are included those that do not produce melanin, like *Fusarium*, *Penicillium*, *Aspergillus*, *Phoma* and those that are black pigmented like *Alternaria*, *Ulocladium*, *Cladosporium* melanin producers. They are fast growing fungi. (ii) The second group includes the so-called black yeasts and meristematic fungi. They are a wide and heterogeneous group of black pigmented fungi that share common characteristics such as the presence of melanins within the cells (swollen cells), hyphae and/or spores. The production of melanin and the meristematic development allow them to survive in stressed environmental conditions like low humidity and high sun irradiation [6]. For this group of fungi is also used the term of rock-inhabiting fungi to underline the exclusive isolation of many of them from rock surface [8]. The black fungi have the capacity to: (i) settle on the rocks surface, (ii) attach firmly to the surface and (iii) penetrate deeper into the rock. The major aesthetic damage, however, may occur when environmental conditions do not force fungi into the crevices, but due to favourable conditions for fungal growth they spread over the rock surface [14].

To preserve constructions from fungal colonisation and to act efficiently against fungal biodeterioration, it is necessary to have a better understanding of biodeterioration mechanisms and its effects on materials properties. Synthetic stone such as concrete, brickwork and mortar show microbial colonisation once their initial highly alkaline pH (>12) falls after a period of weathering (<10) [1]. At the moment, tests to study biodeterioration of building materials exist. Some of them were developed without accelerated weathering of the matrix and so lead to experiment time ranging from 7 to 15 months [46, 56]. On the other hand, accelerated weathering of the matrix may reduce the duration of the experiment by increasing matrix bioreceptivity. Whereas accelerated weathering of the matrix to enhance its bioreceptivity is not truly innovative in biodeterioration study [3, 10, 11, 18, 19, 45, 49], only very few studies [10, 49] focus on the accelerated test for fungal biodeterioration.

An accelerated laboratory test which allows us to compare the growth of three fungal strains and the biodeterioration involved on a cementitious matrix was developed. We paid particular attention to accelerate the matrix weathering so as to not exceed 3 months of experiment. Three fungal strains were selected for the test in order to represent main kind of fungi involved in biodeterioration in natural environment [7, 24, 51, 55, 57, 63, 65]: *Alternaria alternata* to represent a melanin producer *Hyphomycete*, *Exophiala* sp. for yeast like fungi, and *Coniosporium uncinatum* (MC 557) as meristematic fungi. Aesthetical and physical biodeterioration was expected from the development of *Alternaria alternata*, *Exophiala* sp., *Coniosporium uncinatum* with hyphae penetration inside the matrix.

The aim of this paper is to present the accelerated weathering operation which promotes fungal development on hardened cement paste: carbonation and leaching operations are performed to reduce matrix surface pH. The specimens (weathered and non-weathered) are first physico-chemically characterized, and then influence of the matrix weathering on the fungal development is studied.

II. Background

Once the cement based material has been made, leaching and ageing, and more specifically carbonation, start changing the original characteristics of the material. In nature, carbonation and leaching occur intermittently and alternately during the life cycle of the material [62].

II.1. Carbonation

Carbonation is considered the most common chemical reaction influencing the performance of cement based materials in natural environment scenarios [38].

The penetration of gaseous carbon dioxide within partially saturated concrete usually initiates a series of reactions with both ions dissolved in the pore solutions and the hydrated cement paste. The whole process can be summarized as a series of different steps [27, 29, 54]: Gaseous carbon dioxide first penetrates the material. It then dissolves in pore solution mainly as HCO_3^- and CO_3^{2-} . The CO_3^{2-} species then reacts with dissolved calcium to precipitate calcite, CaCO_3 , as well as other CO_2 -based solid phases. The pH drop associated with reactions leads to the dissolution of portlandite. Because carbonation products have higher molar volumes than the parent hydrates Ca(OH)_2 and C–S–H, a decrease of porosity is expected [27, 60].

Progressive carbonation eventually leads to depletion of portlandite and subsequently decalcification of C–S–H to provide new portlandite for the carbonation reaction [27]. This decomposition of C–S–H is reported to increase [27] or decrease [44] porosity.

Since carbonation process in natural environments may be very slow, laboratory experimentation has been conducted under a variety of carbon dioxide levels. Many of the studies used low levels of CO_2 (i.e. 4–5% CO_2) [2, 13]. Accelerated carbonation studies using 100% CO_2 [2, 21, 36, 37, 59]; allow the characterization of carbonation processes in much shorter times.

II.2. Leaching

The leaching of ions (mainly calcium and hydroxide) from the pore solution to the external environment is responsible for the dissolution of portlandite and C–S–H [29]. The leaching of calcium is a coupled dissolution/diffusion process. Leaching by deionized water induces calcium and hydroxide concentration gradients which continuously decrease from the sound zone to the exposed surface of the material. This causes the diffusion of calcium and hydroxide ions from the pore solution to the aggressive solution, and thus lowers the amount of calcium concentration in the pore solution [29]. Loss of calcium leads to the dissolution of portlandite and secondary precipitations of AFm, ettringite and calcite [22, 23, 29].

The degraded zone induced by water exposure is characterized by a decalcification of C–S–H inducing a silicate polymerization. The Ca/Si ratio of the C–S–H gradually decreases between sound and leached zone [29, 34, 39].

Other factors affecting the rate and amount of leaching include the type and amount of constituent investigated, the alkalinity of the matrix, and the chemical properties of the pore water and leachant (pH, ionic strength, etc,...) [62].

Alteration of the chemical behaviour of the matrix resulting from carbonation influences the release of species from the matrix during leaching [25]. Carbonation and leaching are widely investigated together in studies for the use of hazardous wastes in cementitious matrix. Many studies have investigated the effect of carbonation on the leaching of heavy metals [25, 37, 48, 59, 61, 62]. In these studies, the carbonation is mainly performed during the cement hydration. Few focuses on the leaching of carbonated specimens after the curing period, from microstructural point of view. With respect to the effect on pore size distribution, both leaching and carbonation decrease the amount of smaller pores [62].

The research presented here was carried out to evaluate the changes undergone by the cementitious matrix as a result of accelerated weathering. The specific objectives were (i) to characterize the cementitious matrix after each weathering step (carbonation and leaching), and (ii) to evaluate the influence of the matrix accelerated weathering on the fungal growth.

The microstructure of the cementitious matrix is investigated by X-ray diffraction (DRX), Fourier Transform Infrared spectroscopy (FT-IR), Scanning Electron Microscopy with X-ray microanalyses (SEM/EDS).

III. Materials and methods

III.1. Matrix preparation

The study was conducted with ordinary white Portland cement CEM I 52.5 R. The Water/Cement ratio determines the capillary porosity of the cement paste: the higher this ratio, the higher the capillary porosity, but also the higher is the bleeding effect. A higher porosity favours microbial development [31]. The best compromise between bleeding effect and capillary porosity is obtained in our case with a W/C mass ratio of 0.55. Hardened cement paste samples are prisms, $1 \times 2.5 \times 8.5$ cm. Samples were demolded 24 h after production and stored 28 days at 100% relative humidity, at room temperature. Accelerated weathering is performed with carbonation for 48 h, or carbonation (48 h) and leaching operation (28 days) as described by Wiktor *et al.* [64]. Briefly, specimens are exposed to pure CO₂ flow under controlled relative humidity (65% HR) for carbonation. Then, the carbonated specimens are dived into deionized water which is totally renewed in 24 h. Specimens were exposed overnight to UV light for sterilization.

III.2. Analyses

III.2.1. X-ray diffraction (DRX)

The XRD analyses were carried out using Cu K α radiation ($\lambda = 1.5406$ Å) on Bruker D8 advance device. The XRD scans were recorded from 5 to 59° 2 θ with 0.040 step width and 5 s counting time. The analyses were directly performed on the bulk specimen; hence about the first hundred micrometers were analyzed.

III.2.2. Fourier transform infrared spectroscopy (FT-IR)

FT-IR analyses were carried out with Biorad FTS 185 (digilab) equipment using Attenuated Total Reflexion (ATR) unit. The spectra were usually recorded in the range of 4,000–400 cm⁻¹ with 2 cm⁻¹ resolution, and 32 scans were collected each time. The ATR analyses require very small amount of sample (~1 mg), and furthermore no preparation or dilution of the sample is needed. The surface of our specimens was gently scraped and analysed. Hence FT-IR analyses provided information from the first 500 μ m.

III.2.3. Scanning electron microscope equipped with X-ray microanalyses (SEM/EDX)

The fractured surface samples were obtained from the specimens non-weathered, carbonated then leached, and carbonated only. Specimens' cross-sections were also prepared. They were vacuum impregnated with epoxy resin, and polished sections were prepared in the Civil engineering department of Ecole des Mines de Douai.

For the investigation by SEM and EDX, the samples were dried in a desiccator for 24 h, and subsequently gold/palladium coated. Observations were carried out with a JEOL 840 SEM/EDX. Figure 1 shows the different areas analyzed for the EDX analyses.

Each analysis is performed on a surface about 20 \times 200 μ m. Calcium, silicon, and aluminium are assayed.

III.3. Microbiological techniques

III.3.1. Fungal strains and cultural media

Alternaria alternata and *Exophiala* sp. were isolated from a monument in India (unpublished data). *Coniosporium uncinatum* MC557 was isolated from a Carrara marble statue located in the court-yard of the Messina Museum, Italy [9]. These isolated strains were kept in the collection of Department of Microbiological, Genetic and Molecular Sciences in Messina.

Alternaria alternata was cultivated in solid medium (Potato Dextrose Agar—PDA, Oxoid) for 5 days at 26°C. *Exophiala* sp. and *Coniosporium uncinatum* were cultivated in liquid medium (Malt Extract Broth—MEB Oxoid) for 5 days at 26°C.

Nutritive medium used for biodeterioration test is composed of 1 \times concentrated Yeast Nitrogen Broth (YNB) (Difco) plus the addition of Glucose 0.01%.

III.3.2. Fungal units' suspension

Fungal colonies growing in solid medium were scraped from the agar surface and conidia suspended in physiological solution (0.9% of NaCl in distilled water), while the cultures in liquid medium were directly transferred in physiological solution. Each suspension was centrifuged 15 min at 2,095g (3,000 rpm with a Beckman and Coulter Allegra X-12R centrifuge—4750 A rotor). Centrifugation pellets were put in suspension in 1 ml of physiological solution. The number of fungal units was determined through a direct microscopic count in a counting chamber (Bürker chamber, ProSciTech, Australia) and adjusted at a concentration of 8.7×10^5 fungal units/ml in YNB 1× + glucose 0.01% medium.

III.4. Biodeterioration test

III.4.1. Experimental set up

Polyethylene boxes of $9.5 \times 9.5 \times 9.5$ cm³ were used. In order to keep the humidity inside the box, the bottom was covered by vermiculite; paper sheet is disposed on it to avoid the direct contact between specimens and vermiculite. Boxes are autoclaved 15 min at 120°C, then sterile water is added on vermiculite to wet it. Two specimens of each matrix were disposed in each box (Figure 2).

III.4.2. Inoculation

Each specimen was inoculated with 1.5 ml of fungal units' suspension, except controls (only 1.5 ml of sterile medium). Inoculation was performed in duplicate, thus for each strain, 6 specimens were inoculated (2 unweathered, 2 carbonated, 2 carbonated and leached), and placed in 3 different boxes. Boxes were incubated at 26°C.

III.4.3. SEM observation

After 4 weeks of incubation, one specimen of each box was taken and broken in small pieces for SEM observation (Figure 3). Fixation of samples was performed overnight at 4°C in buffered aldehyde fixative 2% (w/v) formaldehyde (P6148, Sigma). Samples were then washed 3 times in 0.01 M phosphate buffer (0.01 M NaH₂PO₄·H₂O, 0.01 M Na₂HPO₄, pH 7.2) for 10 min each. Dehydration was made in graded ethanol series (70, 85, 95% (v/v) and in anhydrous ethanol (100%) for 15 min each). Samples were air dried and then coated with gold for SEM observation (JEOL 840 SEM).

IV. Results and discussion

IV.1. Accelerated weathering

IV.1.1. XRD analyses

The assignments of the XRD peaks to phases were based on 2θ values and the corresponding d-spacing (Figure 4). The peaks corresponding to portlandite appear at 4.900, 3.112, 2.628, 1.927, 1.796, and 1.687 Å. The main peaks corresponding to ettringite are present at 9.671, 5.583, and 3.859 Å. Presence of calcite (CaCO₃) and gypsum can be observed but at a lower extent.

Additional peaks are noticed on the diffractogram of the carbonated specimen compared to non-weathered one. They are attributed to the formation of calcium carbonate polymorphs—aragonite and vaterite—as a result of accelerated carbonation. The decrease in relative intensity of portlandite peaks is due to the dissolution of portlandite to form calcium carbonate. The presence of portlandite can be due to the fact that the depth investigated is larger than the carbonation front. The carbonation operation was performed for 48 h, hence the specimen is only superficially carbonated. The diffractogram of the carbonated and leached specimen exhibits only calcite and aragonite peaks. The disappearance of portlandite and ettringite is a consequence of their dissolution as a result of leaching. Vaterite is the less stable calcium carbonate polymorph, and it is easily transformed into other forms.

IV.1.2. FT-IR analyses

Usually band assignments reported in literature are deducted from spectra performed in diffuse reflectance. In the present work, spectra acquisition is mainly performed in ATR. Band

assignments on ATR spectrum are deducted from spectrum performed first in diffuse reflectance (Table 1).

A shift of about 10–20 wave number units is noted between both spectra (Figure 5). Nevertheless the band assignments for hydrated cement paste are in close agreement with those previously reported [20, 40, 43]. The band at $3,645\text{ cm}^{-1}$ is due to the OH band from Ca(OH)_2 [42]. The water bands appear at approximately $3,440$ (stretching) and $1,660$ (bending) cm^{-1} [41]. The bands at $1,480$ – $1,430$, 875 and 720 cm^{-1} are due to carbonates.

The wide and deep band between $1,480$ and $1,430\text{ cm}^{-1}$ is attributed to ν_3 of CO_3^{2-} and the sharp bands at 875 and 720 cm^{-1} are due to ν_2 and ν_4 vibrations, respectively [41]. The band at 975 cm^{-1} is due to the Si–O asymmetric stretching band (ν_3) from C–S–H [40].

The spectral data changes upon weathering and analyses of these changes can provide valuable information about the reactions involved. The FT-IR data for band assignments are presented in Table 2. The OH band at $3,636\text{ cm}^{-1}$ from portlandite disappears with carbonation. However, XRD analyses show the presence of portlandite on carbonated specimen, but in small amounts. The majority of portlandite is mainly transformed during carbonation operation, and totally during leaching. The small amounts present on the carbonated specimen are probably not detectable with FT-IR, certainly due to smaller penetration depth of analysis on FT-IR compared to XRD.

The FT-IR spectra of weathered specimens (carbonated and leached or carbonated only) show a strong broad band at $1,414$ – $1,440\text{ cm}^{-1}$, which is broader than the calcite band, characteristic of aragonite. According to the intensity it can be assumed that the aragonite overlays the calcite band. New bands appear at $1,080$, 858 , 702 cm^{-1} and are assigned to aragonite formed as a result of accelerated carbonation. The band at 742 cm^{-1} belonging to vaterite, appears only in the carbonated specimen. Other characteristic bands at 860 , 850 , and 710 cm^{-1} may be overlaid by calcite bands. Vaterite is a less stable polymorph of calcium carbonate and thereby transforms into more stable forms calcite and aragonite with leaching operation. The Si–O stretching (ν_3) band now appears at approximately 970 and 980 cm^{-1} for the carbonated and carbonated then leached specimens respectively, while the same band in the non-weathered specimen appears at 959 cm^{-1} . Hence, this shifting to higher frequency is caused by the polymerization of the orthosilicate units (SiO_4^{4-}) during weathering process. The magnitude of this shift is indicative of the degree of polymerization [40].

IV.1.3. SEM observations

Figure 6 shows surface fractures. Platy crystals and fine needles are the distinguishing characteristic shapes of portlandite and ettringite respectively [15]. A thin layer covers progressively the specimen surface as accelerated weathering progresses. This is consistent with the weathering performed: carbonation results in a decrease in the amount of portlandite and the leaching gives rise to the dissolution of portlandite and ettringite. A net decrease in porosity is also observed: pores are progressively filled by CaCO_3 precipitation and become smaller and smaller as the matrix is weathered.

Now, if one examines the following images of polished sections, microstructure alteration is not observed between non-weathered and carbonated specimens (Figure 7a, b). Contrarily, numerous cracks are observed for the carbonated and leached specimen (Figure 7c).

These cracks are distributed along the periphery (Figure 8). This phenomenon may be explained by one of two scenarios: (i) the cracks appear progressively during leaching operation, this could be confirmed by direct observation or (ii) the matrix structure is weakened by the leaching operation, and thereby the cracks are due to the sample preparation. We also notice for this sample the presence of a thin band which seems to be more porous than the rest of specimen (Figure 7c). Its thickness is estimated to be $70\text{ }\mu\text{m}$.

IV.1.4. SEM/EDX analyses

Chemical change was expected from EDX analysis. Figure 9 presents EDX analysis performed on carbonated and leached specimens. Twenty punctual analyses are performed for each area; mean values are plotted on graphs.

We compare concentrations in calcium, silicon, aluminium between analyses performed on the side and in the core. We assume that accelerated weathering has a superficial impact, so the chemical composition in the core is representative of a non-weathered matrix.

No change in chemical composition is observed, there is just a tendency of Ca enrichment and thereby Si depletion of specimen.

IV.1.5. Discussion

Our results show that accelerated carbonation results in calcite, vaterite and aragonite formation (Figure 4; Table 2). This is consistent with literature. The presence of metastable forms (vaterite and aragonite) in the carbonated area seems to be a feature of accelerated carbonation tests [54]. According to Thiery [53], the thermal stability of CaCO_3 produced is lower as the carbonation level increases. This author also showed that the thermodynamically metastable forms of CaCO_3 correspond to the carbonation of C–S–H. The shift of the $\nu_3\text{SiO}_4^{4-}$ from the FT-IR spectrum (Table 2) to higher wave number units suggests also a polymerisation of C–S–H with the carbonation as well as with leaching operations [40].

The carbonation reaction mainly has two effects on the cement matrix and its pore water solution [60]: lowering of the pH of the pore water, and decreasing of the matrix porosity. This is because of the formation of calcite, which is less dense than portlandite, thereby can clog pores more easily. Evolution of total porosity with carbonation is dependent of w/c ratio and cement type.

Generally, the leaching of cement paste gives rise to matrix decalcification due to the dissolution of portlandite and C–S–H decalcification. Haga *et al.* [33] concluded from their work that the major leached constituents of hardened ordinary Portland cement are portlandite and C–S–H gel, and the large pore size, associated with the leaching of portlandite, will significantly affect the diffusion of leached constituents. In the present study leaching occurs after carbonation. The porosity decrease diminishes the leaching of matrix constituents, while the decrease in pH increases leaching. The effect of porosity, however, exceeds that of pH, resulting in a net decrease of leaching due to carbonation [62]. It can be assumed that the dissolution of portlandite and decalcification of C–S–H result in CaCO_3 precipitation in the pore volume. Gervais [26] found that carbonation prior to leaching operation results in a decrease of calcium release compared to the leaching of non-carbonated specimens. The mobility of Ca^{2+} is reduced, and it may precipitate with CO_3^{2-} before, thus not reaching leachate. This could explain EDX analyses and lack of the matrix decalcification observed (Figure 9): most of the dissolved Ca^{2+} precipitates into CaCO_3 before, hence not exiting the matrix with the outcome: enrichment in calcium near the specimen surface.

IV.2. Biodeterioration test

IV.2.1. Observations of non-weathered specimens

Figure 10 presents direct observations of the non-weathered specimens performed after 4 weeks of incubation. No microbial growth is noticed on all exposed specimen. *Coniosporium uncinatum* cells are dark pigmented and taller than those of other fungi inoculated. Hence, this results in the presence of black spots on the specimen. Moreover, when the suspension of cells is inoculated on the non-weathered specimens, it does not penetrate immediately into the matrix. Thus, cells can agglomerate on the surface, resulting in observation of black spots. This spots are, in this case, absolutely not due to the fungal development.

IV.2.2. Observations of carbonated specimens

Figure 11 shows direct observations performed on the carbonated specimens just after inoculation (T0) and after 4 weeks of incubation (T4). No microbial growth is noted on the control and specimens inoculated with *Exophiala* sp. A fungal development is observed on one specimen inoculated with *Coniosporium uncinatum*. It appears after 3 weeks of incubation and looks like small dark area of about 1 cm diameter. Nevertheless, it doesn't seem to be characteristic of *Coniosporium uncinatum* development, but looks more like *Alternaria alternata*. While boxes are handled with lot of care, they remain very close during experiment, so a fungal contamination can easily occur. This should be confirmed with

microscopic observations. Interesting results are obtained with the specimens inoculated with *Alternaria alternata*. Therefore, a fungal development is noticed after 1 week only of incubation for one specimen.

For the second specimen inoculated, it appears after 3 weeks of incubation. The fungus grows until the fourth week. It appears as dark area, scarcely spread on the specimen surface. Hydrated cement paste are heterogeneous material, and there are likely to exist micro-regions that could be prone to preferential fungal colonisation due to *e.g.* differences in the carbonation rate. Differences in specimen porosity would lead to different rates of CO₂-ingress, which in turn would produce varying degree of carbonation. This would result in differences in the pH values in the micro-niches, thus influencing fungal growth [49]. The fungal spreading may also reflect the distribution of fungal cells on the specimen surface after inoculation.

IV.2.3. Observations of the carbonated then leached specimens

Figure 12 presents direct observations performed on the carbonated then leached specimens. Microbial development is noticed on all the specimens after 1 week of incubation only. For the specimens inoculated with *Exophiala* sp. and surprisingly also controls, it appears due to pink coloured spots which seem to grow along the specimen cracks. It is certainly due a bacterial contamination. Moreover, the specimens were inoculated in two series with 1 day of interval: firstly inoculation with *Exophiala* sp., and controls were performed, then the day after with *Alternaria alternata* and *Coniosporium uncinatum*. The bacterial contamination occurred probably during the inoculation of the first series. Nevertheless, this contamination remains localized on restricted area and doesn't spread all over the surface.

Exophiala sp. growth is noticed after the first week on both specimens inoculated. The bacterial contamination doesn't seem to prevent the fungal development. The growth is more pronounced on the periphery area of the specimen surface. No microbial element is observed on the SEM examination of the cross sections.

Alternaria alternata development is observed since the first week of incubation. It is characterized first with mycelial development and then sporulation is observed on the surface. The growth appears more homogeneously than for the carbonated only specimens. SEM observations point out numerous hyphae on the surface (Figure 13a). Crystals, plates like, resulting probably from the sample preparation, are noted from observations of the specimen surface. The cross-section observations show also hyphae but in a more dispersed way (Figure 13b). It may be explained by the fact that fungi can penetrate inside the matrix only by the open porosity, provided the pore diameter is large enough, or through the existing cracks. Therefore, the space available for the fungal growth inside the matrix does not permit extensive development. Nevertheless, the resulting biodeterioration could be more intense inside the exposed material than on the surface.

The development of *Coniosporium uncinatum* is observed since the first week of incubation. It is characterized by the increase of the number and the size of the black spots. Mycelial growth is noted as the incubation time increases. The development seems to occur all over the surface, rather homogeneously. SEM pictures of the surface exhibit the hyphae spread on all over the surface (Figure 13c). Moreover, hyphae are closely linked to the matrix, and even seem to penetrate inside it. Examinations of the cross-section (Figure 13d) clearly show hyphal penetration into the matrix through cracks. It points out the capacity of the fungi to develop not only superficially but also inside the matrix, via structural default, accentuating the physical deterioration.

IV.2.4. Discussion

The results point out that in the present study the microbial growth is promoted with the accelerated weathering of the matrix. It plays a major role. Decrease of surface pH increases considerably matrix bioreceptivity. The microbial colonisation is observed on some carbonated specimens and on all carbonated then leached specimens. Carbonation is the most common chemical reaction influencing cement-based materials in natural environmental scenarios [26, 38]. This is the reason why accelerated weathering of matrix is generally

performed by carbonation [10, 26, 49]. But cement based materials are also exposed to the elements (humidity, acidic rain, snow...) which leads to cement compounds leaching [4]. In our case, carbonation is followed by leaching operation. This allows to obtain a surface pH about 8.8 after 30 days (data not shown), but also to have two ways of weathering involved in natural weathering of materials. To our knowledge, only Escadeillas *et al.* [19] and de Muynck *et al.* [11] performed these two ways of accelerated weathering of the matrix, but giving greater importance to the carbonation step.

Physical weathering of a mineral substrate acts to enhance or accelerate rates of chemical, biomechanical and biochemical weathering and vice versa. The development of cracks, fissures and weathering rinds in rocks accelerates biological weathering by providing a niche that can easily be exploited by opportunistic micro-organisms [5].

The presence of *Alternaria alternata* and *Coniosporium uncinatum*, two melanin producers, was noted on the SEM observations of the cross-sections. Melanin pigmentation of rock-inhabiting fungi confers extra-mechanical strength to the hyphae that are then better able to grow into crevices [17, 30, 52]. Increased penetration augments contact of fungal hyphae and their metabolites with the rock, amplifying their geochemical influence on the mineral substrate. Expansion of hyphal growth into deeper rock layers also helps to protect the cells from the UV radiation [30]. One characteristic of meristematic fungi is the ability to form filamentous hyphae that develop from clump-like colonies to penetrate deep into rocks thus protecting themselves from environmental stresses. In this sense, the visible portion of melanized micro colonial fungi is like the tip-of-the-iceberg, because the hyphae can rapidly penetrate several mm to cm into hard rocks in search of more protected environments [30]. The fungi form explorative hyphae to seek nutrients on the surface, or mycelium during the growth. Under natural conditions, nutrient availability, even if small, enhances the chance of fungal units (single cells, conidia, or hyphae fragments) surviving and starting the colonization [55, 56].

V. Conclusion

Results showed that there was no microbial development when the cementitious matrix was not weathered. Microbial growth was noted on and inside the carbonated then leached specimens.

Relating to the matrix weathering; combined process led to the maximum microbial growth. This process was composed by a carbonation operation followed by the leaching of the matrix. This resulted in modification of the matrix microstructure and above all in the surface pH decrease. Carbonation of hardened cement paste led to a surface composition nearer of a limestone than that of a cementitious matrix.

The accelerated laboratory test developed in this study permits to obtain a rapid fungal development on cement specimens. Three months of experiments only are needed to obtain first results, which is rather shorter than other test developed to date to study fungal biodeterioration. Results are mainly related to aesthetical biodeterioration. Moreover, fungal strains presenting various growth patterns were used, and so may lead to different biodeterioration mechanisms in long term exposition experiment. These results will be completed by the quantification and estimation of the extent of the fungal growth.

This test could be used to compare the bioreceptivity of various cement based materials, leading to results in 3 months only.

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Figures

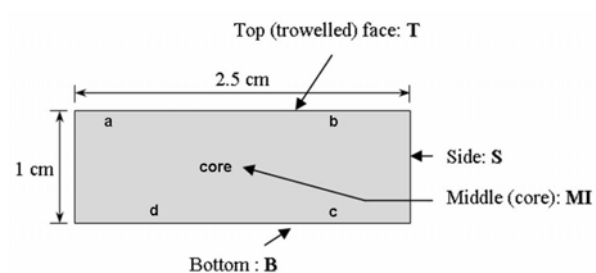


Figure 1: Localization of EDX analyzed areas.

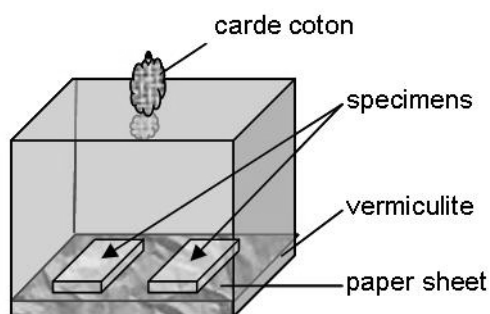


Figure 2: Experimental set up for the biodeterioration test.

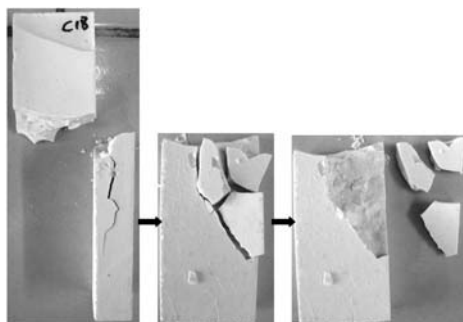


Figure 3: Example of a specimen broken to carry out SEM observations.

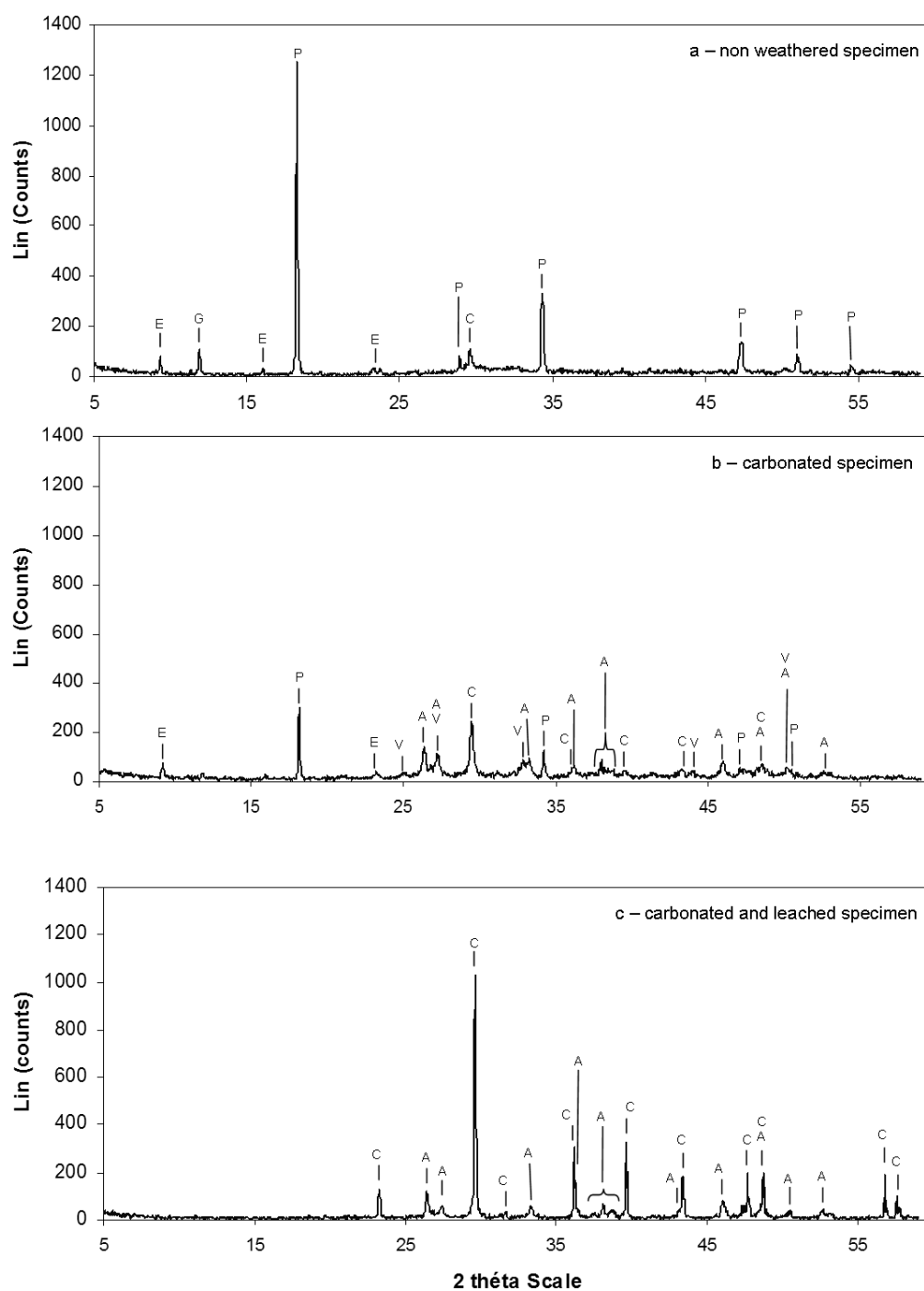


Figure 4: X-ray diffractogram of specimens before (a) and after (b), (c) accelerated weathering of the matrix – E=ettringite, G=gypsum, P=portlandite, C=calcite, A=aragonite, V=vaterite.

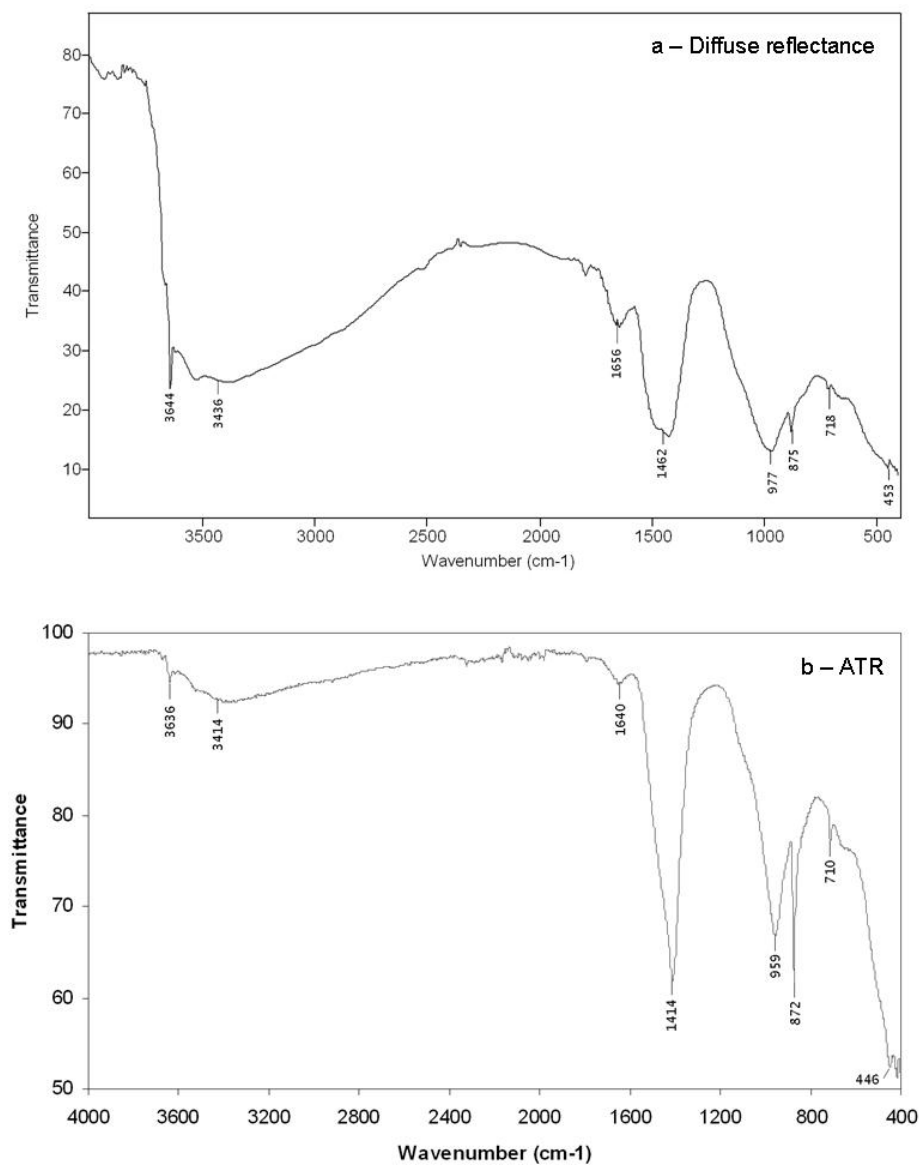


Figure 5: FT-IR spectra of hydrated cement paste at 28 days of curing at 100%RH – (a) diffuse reflectance, (b) ATR.

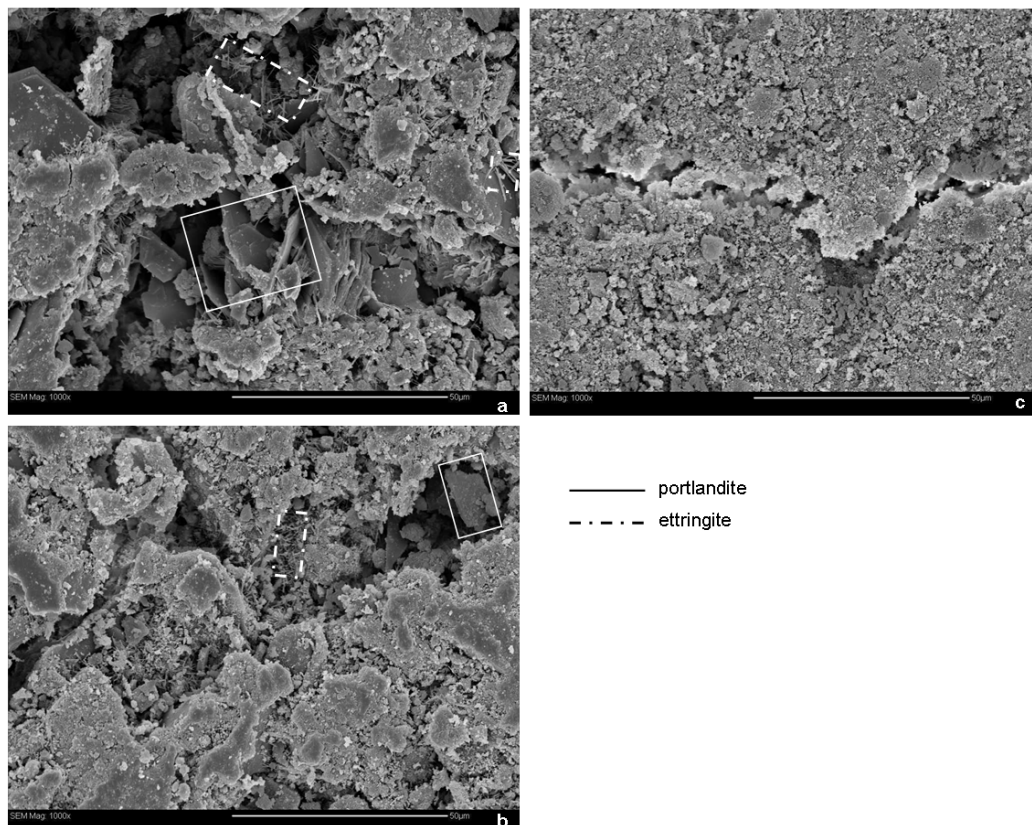


Figure 6: SEM images of surface fractures of specimens – (a) non weathered, (b) carbonated, (c) carbonated then leached.

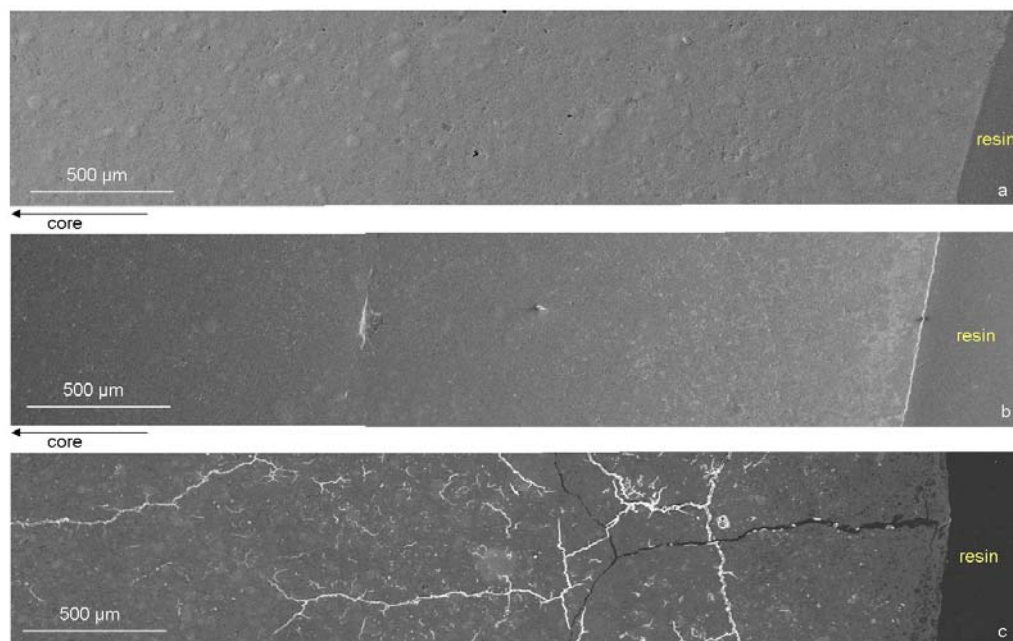


Figure 7: SEM images of polished sections for specimens (a) non weathered, (b) carbonated, (c) carbonated then leached.

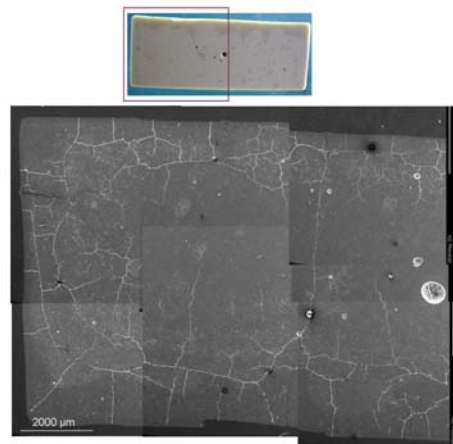


Figure 8: Global SEM images of polished sections of a carbonated then leached specimen.

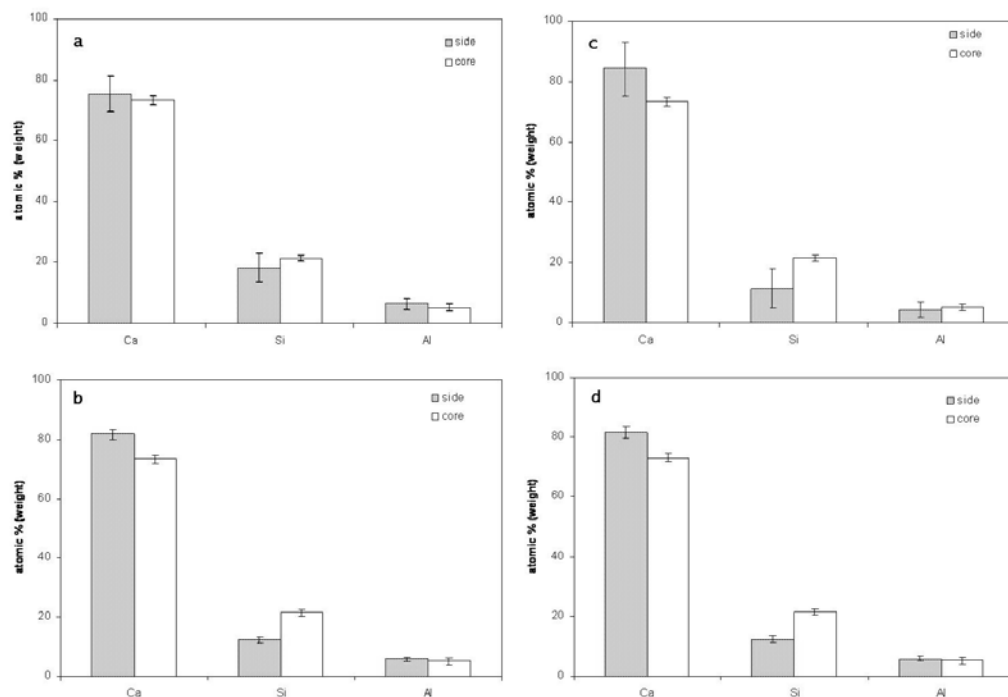


Figure 9: SEM/EDX analyses of polished sections for carbonated then leached specimen – a, b, c, d correspond to the analyzed areas named in section III.2.3.

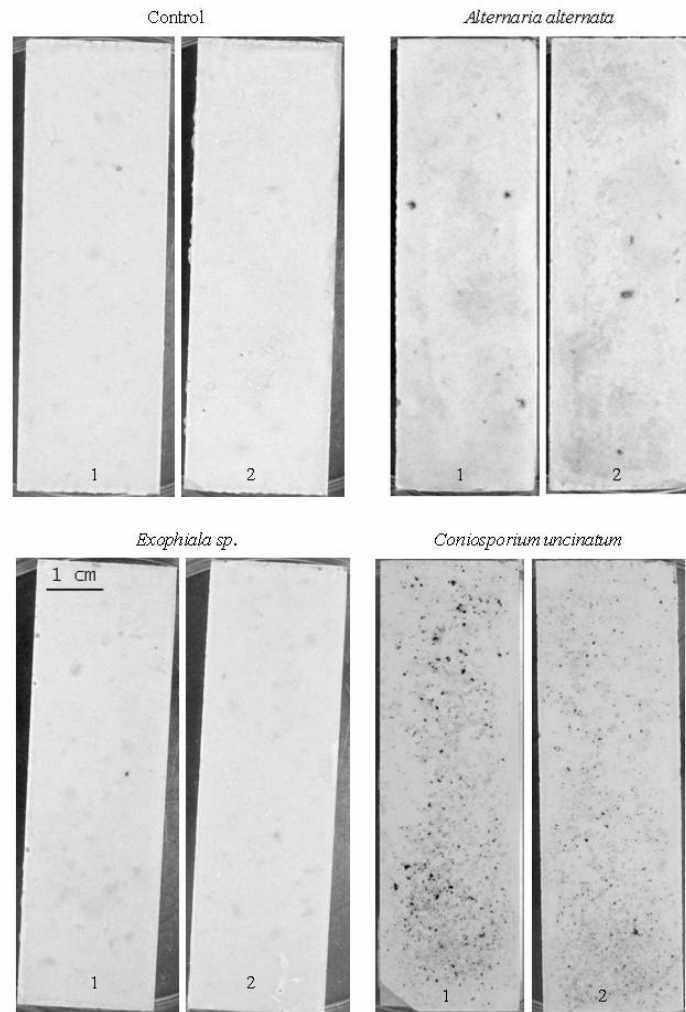


Figure 10: Direct observations of the non weathered specimens after 4 weeks of incubation.

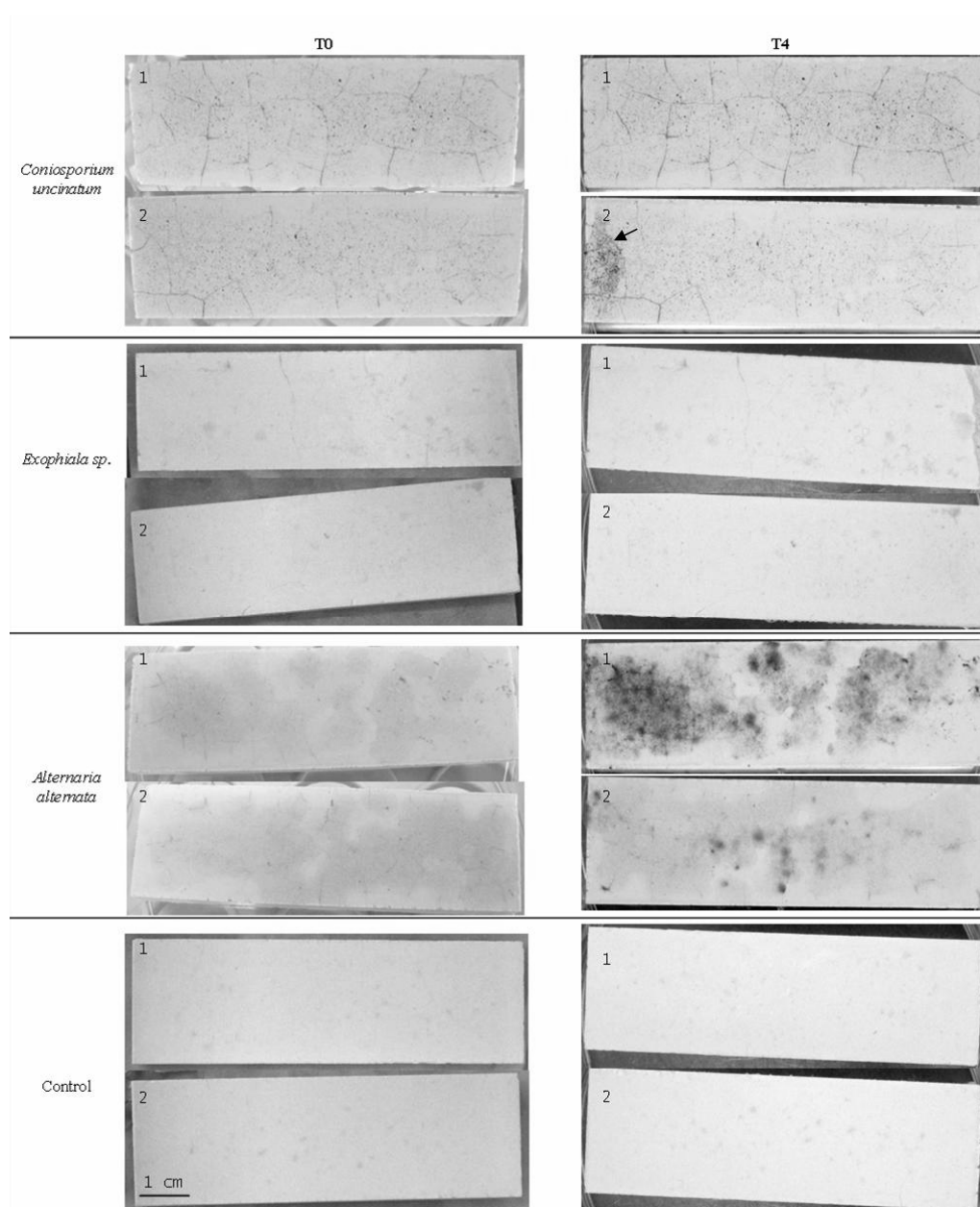


Figure 11: Direct observations of the carbonated specimens – arrow indicates contamination.

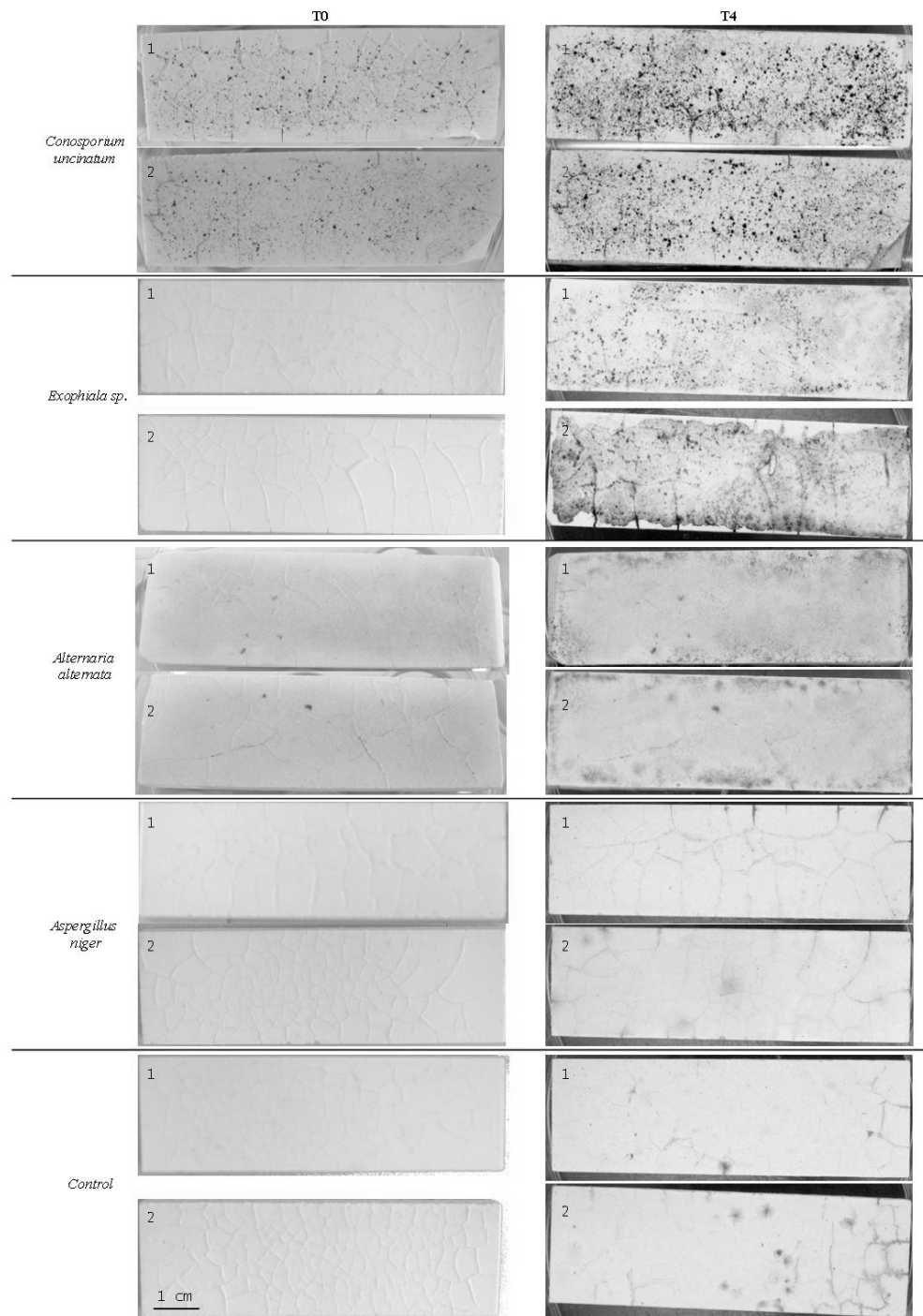


Figure 12: Direct observations of the carbonated then leached specimens.

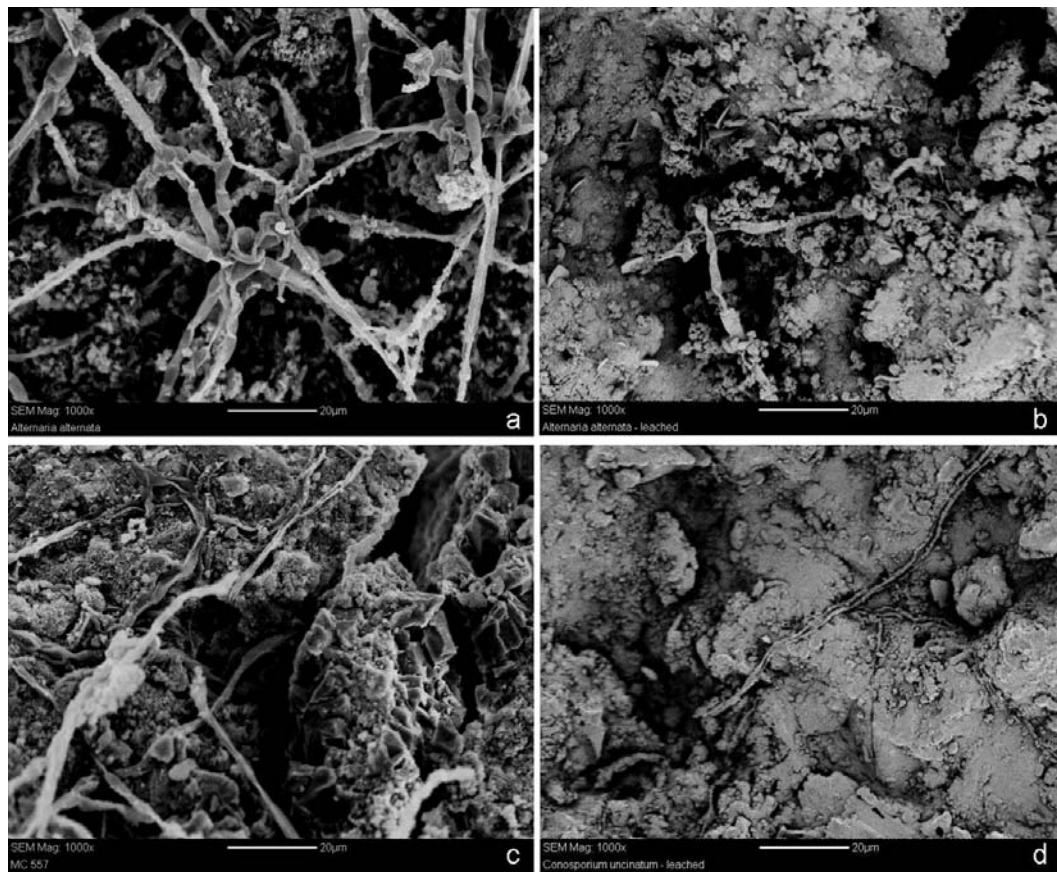


Figure 13: SEM observations of the carbonated then leached specimens – Surface (a) and cross-section (b) of specimens inoculated with *Alternaria alternata* - Surface (c) and cross-section (d) of specimen inoculated with *Coniosporium uncinatum*.

Tables

Table 1: FT-IR characterization of hydrated cement paste.

Band assignments	Hydrated cement ^a 1 month (cm ⁻¹)	Hydrated cement ^b after mixing (cm ⁻¹)	Hydrated cement ^c paste (cm ⁻¹)	Hydrated cement ^d paste (cm ⁻¹)
$\nu_3 \text{SiO}_4^{4-}$	980 s, b	970 s, b	977 s, b	959 s
$\nu_4 \text{SiO}_4^{4-}$	536 w, sh	—	—	—
$\nu_2 \text{SiO}_4^{4-}$	467 s	—	453 w	446 w
$\nu_3 \text{SiO}_4^{2-}$	1105 w, sh	1100	1134 vw, sh	1113 vw, sh
	1155 w, sh	—	—	—
$\nu_4 \text{SiO}_4^{2-}$	667 w	610	—	—
$\nu_1 + \nu_3 \text{H}_2\text{O}$	3325	—	—	—
	3450 s, b	3415 s, b	3436 s, b	3414 m, b
$\nu_2 \text{H}_2\text{O}$	1630 w	1638 m	1656 m	1640 w
VOH^-	3645 sr, sh	3640 sr	3644 sr, sh	3636 sr, sh
$\nu_3 \text{CO}_3^{2-}$	1425 1497 s, b	1421 s, b	1462 s, b	1414 s
$\nu_2 \text{CO}_3^{2-}$	876 m, sr	874 m, sr	875 m, sr	872 s, sr
$\nu_4 \text{CO}_3^{2-}$	732 w	713 w	718 w	710 w

b Broad, s strong, sh shoulder, sr sharp, m medium, vw very weak, w weak

ν_1 symmetric stretching, ν_2 symmetric bending, ν_3 asymmetric stretching, ν_4 asymmetric bending

^aMollah et al. [40, 43]

^bFarcas and Touze [20]

^cPresent study diffuse reflectance

^dPresent study ATR

Table 2: FT-IR characterization before and after accelerated weathering [12, 20, 28, 50].

Band assignments	Calcium carbonate ^a CaCO ₃ (cm ⁻¹)	Non-weathered ^b specimen (cm ⁻¹)	Carbonated ^c specimen (cm ⁻¹)	Carbonated and leached specimen ^d (cm ⁻¹)
ν_3 SiO ₄ ⁴⁻	970 s, b	959 s	970 s	970 s
ν_2 SiO ₄ ⁴⁻	—	446 w	—	—
ν_3 SiO ₄ ²⁻	1100	1113 vw, sh	—	—
ν OH ⁻	3640 sr	3636 sh	—	—
ν_3 CO ₃ ²⁻ calcite	1421 s, b	1414 s	1414 s	1414 s
ν_3 CO ₃ ²⁻ aragonite	1400–1500 s, b	—	1414–1440 s, b	1414–1440 s, b
ν_1 CO ₃ ²⁻ vaterite	1080 w	—	—	ν_1 CO ₃ ²⁻ aragonite
1083	—	1080 w, sh	1080 w, sh	ν_2 CO ₃ ²⁻ calcite
874 m, sr	875 s, sr	876 s, sh	876 s, sh	—
ν_2 CO ₃ ²⁻ vaterite	860 s, sr	—	?	?
850 m, sr	—	?	?	ν_2 CO ₃ ²⁻ aragonite
856 m	—	858 s, sh	858 s, sh	ν_4 CO ₃ ²⁻ calcite
713 w	710 w	714 m	714 m	ν_4 CO ₃ ²⁻ vaterite
740 w	—	742 w, sh	—	710 m

b broad, s strong, sh shoulder, sr sharp, m medium, vw very weak, w weak

ν_1 symmetric stretching, ν_2 symmetric bending, ν_3 asymmetric stretching, ν_4 asymmetric bending, ? certainly covered by another band

^aSimspon [50], Ghosh [28], Farcas and Touzé [20], and Devarajan et al. [12]

^bPresent study—non-weathered specimen

^cPresent study—carbonated specimen

^dPresent study—carbonated and leached specimen